

TIE2-expressing macrophages limit the therapeutic efficacy of the vascular-disrupting agent combretastatin A4 phosphate in mice

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Vascular-disrupting agents (VDAs) such as combretastatin A4 phosphate (CA4P) selectively disrupt blood vessels in tumors and induce tumor necrosis. However, tumors rapidly repopulate after treatment with such compounds. Here, we show that CA4P-induced vessel narrowing, hypoxia, and hemorrhagic necrosis in murine mammary tumors were accompanied by elevated tumor levels of the chemokine CXCL12 and infiltration by proangiogenic TIE2-expressing macrophages (TEMs). Inhibiting TEM recruitment to CA4P-treated tumors either by interfering pharmacologically with the CXCL12/CXCR4 axis or by genetically depleting TEMs in tumor-bearing mice markedly increased the efficacy of CA4P treatment. These data suggest that TEMs limit VDA-induced tumor injury and represent a potential target for improving the clinical efficacy of VDA-based therapies.

Introduction

Vascular-disrupting agents (VDAs) selectively target the tumor vasculature, resulting in a rapid and severe impairment of blood flow and secondary tumor cell death (1, 2). The lead VDA, combretastatin A4 phosphate (CA4P), is active in transplanted and spontaneous murine tumors as well as in human tumors (1, 3, 4). However, viable blood vessels remaining at the tumor periphery provide a focus for regrowth and revascularization of the damaged tumor, representing treatment resistance and limiting the long-term efficacy of CA4P (1, 5).

Tumor-associated macrophages (TAMs) stimulate tumor angiogenesis (6), and high TAM numbers correlate with a poor prognosis in cancer patients (7). A highly proangiogenic TAM subset, the TIE2-expressing macrophages (TEMs), are present in a variety of mouse and human tumors (8–10) and physically interact with newly forming tumor blood vessels (8, 9) and express high levels of several proangiogenic factors (11, 12). Increased TEM infiltration occurs in mouse glioblastoma following high-dose radiation therapy and is associated with tumor revascularization and relapse (13). TEMs have also been implicated in the regrowth of subcutaneous breast and lung carcinomas after local irradiation (14). In the current study, we show that TEMs limit the therapeutic efficacy of CA4P in tumor-bearing mice.

Results and Discussion

Twenty-four hours after a single injection of CA4P, mammary tumors in MMTV-PyMT mice were significantly more hypoxic/ necrotic than those in controls (P < 0.05; Figure 1, A and B) and

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contained CD31⁺ blood vessels with smaller lumens (Figure 1C). This is consistent with the vascular-disrupting effects of CA4P reported in various mammary tumor models (4, 15).

The number of F4/80⁺ TAMs was also significantly increased in CA4P-treated tumors (Figure 1, D and E). F4/80, TIE2, and CD31 immunofluorescence staining of tumors showed that infiltration of CD31⁻F4/80⁺TIE2⁺ TEMs significantly increased 24 hours after CA4P injection (Figure 1, D and E), while the number of F4/80⁺TIE2⁻ TAMs was not significantly altered (data not shown). These data were confirmed by flow cytometric analysis of dispersed tumors (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI44562DS1). Virtually all F4/80⁺TIE2⁺ TEMs expressed the proangiogenic enzyme MMP9 in both untreated and treated tumors, whereas far fewer TIE2-F4/80⁺ TAMs were MMP9⁺ (Supplemental Figure 2, A and B). CA4P treatment also significantly increased tumor infiltration by MMP9-expressing TEMs in N202 mammary tumors grown subcutaneously in syngeneic mice (Supplemental Figure 2, C and D).

Tumor expression of CXCL12 (SDF1) by both tumor and stromal cells (identified on the basis of cell morphology) increased 24 hours after CA4P treatment (Figure 2A and Supplemental Figure 3) and was consistent with the hypoxic upregulation of CXCL12 (16) and raised plasma levels of CXCL12 seen in CA4P-treated patients (17). Ninety percent of tumor TEMs and 60% of TIE2⁻ TAMs expressed the CXCL12 receptor, CXCR4, in MMTV-PyMT tumors (Figure 2B). CXCR4 expression per cell was significantly higher in TEMs than in TIE2⁻ TAMs and was augmented further by CA4P treatment (Figure 2C). We then used the CXCR4 inhibitor, AMD-3100, to investigate the role of the CXCL12/CXCR4 axis in TEM recruitment to tumors after CA4P. This significantly reduced TEM but not TIE2⁻ TAM recruitment to CA4P-treated N202 tumors (Figure 2D). Of note, AMD-3100 plus CA4P treatment elicited a significant increase in tumor necrosis compared with CA4P treatment alone (Figure 2D).

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In order to study the effects of combined CXCR4 blockade and CA4P treatment on tumor responses to CA4P, we used the subcutaneous N202 mammary tumor model, which allows more reliable tumor measurements and monitoring of tumor volume than the MMTV-PyMT model. AMD-3100 alone had no effect on N202 tumor growth (Figure 2E), consistent with previous studies in other tumor models (14). The combined drug treatment (AMD-3100 and CA4P) induced a significant inhibition of N202 tumor growth compared with CA4P alone (Figure 2E).

Figure 1

CA4P induced vascular damage, increased tumor hypoxia, and increased TEM numbers in MMTV-PyMT mammary tumors. (A) H&E staining shows a marked increase in tumor necrosis (N) 24 hours after 50 mg/kg CA4P. (B) Staining with anti-PIMO shows increased hypoxia (red). (C) Tumor sections stained with anti-CD31 antibodies show reduced lumen of blood vessels (white) 24 hours after 50 mg/kg CA4P. (D) Tumors stained with DAPI (to show cell nuclei; blue), anti-F4/80 (green), and anti-TIE2 (red). Yellow arrows denote F4/80+TIE2+ TEMs; F4/80-TIE2+ structures represent blood vessels. (E) Number of all F4/80⁺ cells/field and F4/80⁺TIE2⁺ TEMs/field (by immunofluorescent staining; see D above) and proportion (%) of F4/80+TIE2- and F4/80+TIE2+ cells (TEMs) in viable cells obtained from dispersed tumors (by flow cytometric analysis; see Supplemental Figure 1). Scale bars: 50 µm. **P* < 0.05, ***P* < 0.01, ****P* < 0.001, compared with relevant control group. NS, no statistical difference (P > 0.05).

These results are consistent with the necrosis data obtained in MMTV-PyMT tumors (Figure 2D).

Because CXCR4 is not only expressed by TEMs in tumors, we also used a conditional suicide genebased strategy for specific TEM elimination (8). This involved transplanting BM cells from Tie2-HSV-thymidine kinase transgenic mice into syngeneic, wildtype mice. Six weeks after BM transplant, ganciclovir (GCV) was used to specifically eliminate TEMs in mice carrying established N202 tumors. CA4P induced an increase in necrosis (Figure 3C) and vessel narrowing (Supplemental Figure 4A) in the N202 mammary tumors of transplanted mice. Coadministration of GCV to deplete TEMs resulted in significant TEM depletion in CA4P-injected tumors (Figure 3, A and B) accompanied by a striking increase in tumor necrosis in CA4P-treated tumors compared with non-TEMdepleted CA4P-treated tumors (Figure 3C). Tumor necrosis was most marked at 24 hours after CA4P in the TEM-depleted mice, indicating that TEMs are protective at early stages after treatment (Figure 3C). Interestingly, the number of tumor-infiltrating F4/80-Gr1⁺ neutrophils increased significantly at 24 hours after CA4P treatment and increased further in TEM-depleted mice that received CA4P (Supplemental Figure 4B). However, very few F4/80-Gr1+ neutrophils were seen at 72 hours in any of the treatment groups (Supplemental Figure 4B). Contrary to CXCR4 blockade (Figure 2D), GCV completely abolished TEM recruitment to tumors (Figure 3B) and - in agreement with previous findings (8) - markedly inhibited tumor growth (Supplemental Figure 5). It was there-

fore not possible to assess the additive effects of CA4P treatment and TEM elimination on tumor growth in this model.

Taken together, our results show that TEMs are rapidly recruited into CA4P-treated mammary tumors, where they have a protective effect against vascular damage, necrosis induction, and growth retardation. At present, it is not possible to distinguish whether TEMs are acting to protect against the damaging cascade of events that lead to CA4P-induced tumor necrosis or whether they are promoting rapid revascularization and tumor



Figure 2

Inhibition of tumor recruitment of CXCR4⁺ TEMs increases the therapeutic efficacy of CA4P. (**A**) CXCL12 expression, determined by immunofluorescence staining (see Supplemental Figure 3), was increased in MMTV-PyMT tumors 24 hours after injection with 50 mg/kg CA4P. (**B** and **C**) Flow cytometric analysis of dispersed MMTV-PyMT tumors shows that (**B**) the majority of F4/80⁺TIE2⁺ TEMs express CXCR4 and the MFI for CXCR4 is higher for TEMs than TIE2⁻ TAMs and (**C**) CXCR4 expression/TEM is upregulated after CA4P treatment. (**D**) The CXCR4 inhibitor, AMD-3100, inhibits recruitment of TEMs, but not TIE2⁻ TAMs, to MMTV-PyMT tumors and increases CA4P-induced tumor necrosis. n > 5 mice per group. *P < 0.05; **P < 0.01. (**E**) Twice daily i.p. injections of 5 mg/kg AMD-3100 (for the period indicated) enhanced the effects of 3 daily i.p. injections of 50 mg/kg CA4P on the growth of N202 tumors. Statistical analysis in **E** employed the nonlinear mixed effects model described in the Supplemental Methods.

repopulation of necrotic regions. Considering the proangiogenic roles of TEMs in tumors (8, 9, 12, 18) and our finding that TEMs avidly express MMP9, we speculate that a reparative role is highly likely. Either way, our data indicate that targeting these cells is a promising approach to improving the efficacy of a VDA like CA4P.

Endothelial progenitor cells (EPCs) were previously shown to play an important role in tumor recovery following VDA treatment (17, 19). Shaked et al. (19) depleted EPCs using DC101, a mouse monoclonal antibody to VEGFR-2. When administered to tumor-bearing mice along with a CA4P derivative, OXi-4503 (CA1P), DC101 decreased circulating EPC numbers and tumor blood flow and increased tumor necrosis, compared with OXi-4503 alone. However, DC101 would also have blocked VEGFR-2 signaling in the viable tumor blood vessels and therefore had direct effects on tumor angiogenesis that were independent of EPCs (18). Because BM-derived EPCs are very rare cells in mouse blood and tumors (8, 9, 14, 20–22), it is likely that other BMderived cell types, such as monocyte-lineage cells, are primarily involved in regulating tumor responses following VDA treatment. Although our findings do not eliminate a potential contribution of EPCs to tumor revascularization/recovery after CA4P, they indicate that TEMs play a crucial role in this process.

Interestingly, TEM depletion stimulated an increase in neutrophil infiltration into tumors — even in the absence of subsequent treatment with CA4P, suggesting that TEMs may inhibit tumor infiltration by Gr1⁺F4/80⁻ neutrophils (23). CA4P treatment alone also increased neutrophil infiltration, possibly in response to tumor necrosis, and TEM depletion, in combination with CA4P, increased it even further. It is noteworthy that tumor Gr1⁺ neutrophils had significantly dropped by 72 hours, illustrating that any participation in CA4P-induced changes in tumors was time dependent. Clearly, the enhanced recruitment of neutrophils at 24 hours after treatment failed to protect tumors. Rather, neutrophils may have contributed to the increased level of necrosis present by exhibiting a cytotoxic phenotype (24). Interestingly, levels of neutrophil-derived myeloperoxidase (MPO), indicative of such a phenotype, are associated with CA4P-induced vascular injury in tumors (25).

In summary, our data show that TEMs are rapidly recruited to CA4P-treated mammary tumors and protect against the vascular damaging effects of this archetypal VDA. This suggests that targeting TEMs could increase the long-term efficacy of CA4P.

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Saline

CA4P

GCV

GCV + CA4P



Figure 3

Conditional depletion of TEMs increases the efficacy of CA4P treatment. (**A**) N202 tumors excised 24 hours after the last of 3 daily i.p. injections of CA4P or saline in the presence or absence of GCV to deplete TEMs were stained with DAPI (blue), anti-F4/80 (green), anti-TIE2 (red), and anti-MMP9 (white). TEMs are highlighted with yellow arrows; non-TAMs that express MMP9 are shown with white arrows. Scale bar: 50 μ m. (**B** and **C**) Changes in TEM numbers and tumor necrotic area 24 hours and 72 hours after last CA4P injection. *n* = 6 mice per group. **P* < 0.05.

Methods

See Supplemental Methods for details of methods not outlined below. All animal studies were approved by the University of Sheffield Ethics Committee/UK Home Office regulations and the Animal Care and Use Committee of the Fondazione San Raffaele del Monte Tabor, Milan, Italy.

CA4P studies. Female FVB/MMTV-PyMT mice (10–12 weeks old) were given a single i.p. injection of 50 mg/kg CA4P or saline and culled 24 hours later. Mice were also injected i.v. with 60 mg/kg pimonidazole hydrochloride (PIMO; a nitroimidazole compound used for detecting hypoxia in tumor sections) 1 hour prior to culling. Excised tumors were bisected and half frozen in OCT compound and half fixed in 10% formalin (and then processed into paraffin wax). Some tumors were excised and enzymatically digested for FACS analysis. Mice bearing N202 mammary tumors were administered 50 mg/kg CA4P for 3 consecutive days and tumors excised 24 or 72 hours after the final dose.

AMD-3100/CA4P studies: MMTV-PyMT tumor model. MMTV-PyMT mice were injected i.p. twice daily with 5 mg/kg AMD-3100 (Sigma-Aldrich), a specific CXCR4 inhibitor. On the third day, mice were given a single i.p. injection of 50 mg/kg CA4P or saline (PBS) 2 hours after the first dose of AMD-3100. 24 hours after the last CA4P injection, mice received 1 further injection of AMD-3100 and were culled 2 hours later.

AMD-3100/CA4P studies: N202 tumor model. N202-bearing mice were injected s.c. at day 0 and 5 mg/kg AMD-3100 or saline administered twice daily i.p. starting at day 7 (when tumors reached approximately 150 mm³) and ending on day 16 when mice were killed and tumors removed. Daily

i.p. injections of CA4P (50 mg/kg) or saline started at day 9 and ended on day 13. Tumor size was determined using callipers.

TEM conditional depletion studies. Six-week-old Tie2-HSV-tk transgenic mice (8) were sacrificed with CO_2 and their BM harvested by flushing the femurs and the tibias. BM cells (10⁷/mouse) were infused into the tail veins of 40 lethally irradiated (975 cGy) 6-week-old FVB mice. Six weeks after transplant, we confirmed hematopoietic reconstitution of recipient mice by quantitative PCR (qPCR) analysis of vector sequences in blood leukocytes, as described (8, 9). N202 mammary tumor cells (5 × 10⁶) were then injected s.c. into the flanks of BM-transplanted mice (day 0). Thirteen days (day 13) after N202 tumor challenge (average tumor volume: 700 mm³), transplanted mice were treated with GCV (100 mg/kg) daily for 11 days or remained untreated. Tumor size was determined using callipers. Nineteen days (day 19) after N202 tumor cell inoculation, GCV-injected and non-GCV-injected mice were randomly allocated to receive CA4P (50 mg/kg) or saline vehicle for 3 consecutive days; they were culled 24 or 72 hours later and tumors excised and processed as above.

Statistics. In most cases, multiple comparison of groups was performed by ANOVA followed by the Tukey-Kramer honest significance difference test (GraphPad Software Inc.). See figure legends for any alternative statistical tests used. The non-linear mixed effects (NLME) model used for the tumor growth curves in Figure 2E was implemented using the R statistical software, version 2.9.1 (The R Project for Statistical Computing; http://www. r-project.org/) (see Supplemental Methods). All data represent mean values ± SEM; P values of less than 0.05 were considered to be significant.



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